

# Dealing with indetermination in biochemical networks

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## ABSTRACT

A main aspect in computational modelling of biological systems is the determination of model structure and model parameters. Due to economical and technical reasons, only part of these details are well characterized, while the rest are unknown. To deal with this difficulty, many reverse engineering and parameter estimation methods have been proposed in the literature, however these methods often need an amount of experimental data not always available.

In this paper we propose an alternative approach, which overcomes model indetermination solving an Optimization Problem (OP) with an objective function that, similarly to Flux Balance Analysis, is derived from an empirical biological knowledge and does not require large amounts of data. The system behaviour is described by a set of Ordinary Differential Equations (ODE). Model indetermination is resolved selecting time-varying coefficients that maximize/minimize the objective function at each ODE integration step. Moreover, to facilitate the modelling phase we provide a graphical formalism, based on Petri Nets, which can be used to derive the corresponding ODEs and OP. Finally, the approach is illustrated on a case study focused on cancer metabolism.

## CCS Concepts

•**Computing methodologies** → **Modeling methodologies**; *Uncertainty quantification*; •**Applied computing** → Systems biology; •**Software and its engineering** → Petri nets;

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## Keywords

indetermination, Ordinary Differential Equation, optimization problems, Petri net.

## 1. INTRODUCTION

The inherent complexity of biological systems makes the study of their dynamics and behaviours a difficult issue. Within this context, computational modelling can help biologists/clinicians to identify which molecules and interactions have a crucial role on the global behaviour of the system. Many modelling approaches have been proposed in the literature. An appropriate choice among them depends on the specific biological question being addressed and on the available data. An extensive review of the most successful modelling approaches used for metabolic, and more in general for any biochemical network, is reported in [4], where the authors divide these approaches into three classes (i.e. *interaction-based*, *constraint-based* and *mechanism-based* approaches) depending on the used level of abstraction. For instance, *interaction-based* approaches do not consider any quantitative aspect, so that they allow only a topological analysis of the system. Instead, *constraint-based* approaches rely on the assumption of steady state and typically exploit Flux Balance Analysis (FBA) and its extensions [17] to study the system at equilibrium. *Mechanism-based* approaches describe interactions with high level of detail and are the most likely candidates to provide a complete understanding of biochemical dynamics.

In order to build a mechanism-based the modeller has to specify all its characteristics, like, for instance, the structure of the system, the mechanism of the events that produce changes in variables and the mathematical expressions used to represent them, with all their parameters defined. This considerable amount of preliminary information requires the modeller to acquire an extensive knowledge of the biological phenomena and to formulate all the assumptions on which the model relies. In this article we will focus on a particular type of biochemical systems, namely metabolic systems, and on modelling these systems with Ordinary Differential Equations (ODEs). Due to economical and technical reasons, the complete information needed to build the set of

ODEs is rarely available, causing the modeller to face situations of indetermination. In this study we will consider in particular those metabolic models in which part of the reactions are fully characterized while the rest are not. Two groups of reactions are thus defined: determined reactions, having well characterized structure, mathematical expression and parametrization, and undetermined reactions, described just by their stoichiometry.

Issues of indetermination have been already deeply studied in literature and solutions have been proposed at different levels, depending on the type of missing information. Techniques of reverse engineering (RE) and Parameter Estimation (PE), for instance, overcome this indetermination by means of available experimental data and optimization algorithms [4, 14, 8].

In this article we propose an alternative approach to deal with situations in which insufficient experimental data hamper the application of PE and RE methods. To overcome the lack of information concerning undetermined reactions we chose to exploit an empirical biological knowledge that helps us formulate some modelling assumptions and define an optimization problem (OP).

Since the direct definition of the ODEs and of the OP may be difficult and may require advanced modelling skills, we facilitate this process providing a high level graphical formalism based on Petri Nets (PNs). The paper is organized as follows. Section 2 gives a general outlook on other methods currently used for similar purposes, while Section 3 provides the necessary background on PNs and OPs. In Section 4 our approach is introduced: a formal definition of our PN extension and how to automatically translate a model described through such a high level formalism into the underlying ODE system and OP are provided. In Section 5 some experimental results are shown, and our method is compared to others present in literature. In Section 6 we conclude presenting some future works.

## 2. STATE OF THE ART

As mentioned in the introduction, situations of inadequate biochemical characterization are traditionally managed with **Reverse Engineering** (RE) or **Parameter Estimation** (PE) computational techniques. When the topology of the network is not known, several techniques of RE can be used. Besides topology, the parametrization of the model represents at present the main area of lacking information. The parameters required for a kinetic model include both the kinetic parameters and the initial conditions. Many parameters have been experimentally calculated with biochemical assays, and have been reported in literature, but many others are still unavailable. As some authors have pointed out, if we consider the parametrization of the mathematical expressions of enzyme activity: “for the foreseeable future, full availability of *true* rate equations for all enzymes is certainly an illusion” [3]. To cope with this situation, many PE techniques have proposed different approaches that aim to automatically infer model parameters from experimental data. In general, the inputs of PE methods include network topology, initial conditions and discrete-time time series of some biochemical species present in the model. When parameters and network topology are unknown, PE has to be embedded in the solution of a Reverse Engineering problem [4]. Instead, if parameters and initial conditions are unavailable, a joint state-kinetic constants estimation (see, e.g., [12]) has to

be performed. PE performs a global optimization of a fitness function which evaluates the distance between the simulated solution and the experimental data points. Several alternative optimization algorithms can be found in literature, each adopting a specific approach to escape from local minima and to limit the computational complexity. The difficulty of PE lies in the availability of experimental data, which in many real cases are very scarce. In this paper we present an approach that can be used in situations where both part of the network structure and part of the parametrization are unknown, representing an alternative to a PE embedded in a RE process. Differently from these techniques, our method tries to escape a strict dependency from experimental data using an OP that relies on biological empirical knowledge. This OP exploits an objective function that is conceived to represent a specific biological behaviour or phenotype, as it will be discussed in Section 5.

The use of an objective function with a biological meaning has already been incorporated in **Flux Balance Analysis** and **Cybernetic Modelling** (CM). In most FBA studies the objective function describes an objective that we believe is physiologically relevant for the cell, like the maximisation of biomass. This is used to predict a physiological behaviour of the cell. The idea that a system, like a cell, seeks a biological objective is justified acknowledging the apparent goal-directedness of nature. Other computational approaches, like cybernetic modelling [21], have taken advantage of this concept, which has been formally defined with the word teleonomy. A teleonomic view of nature indeed considers that the genetic programs of all living organisms has been shaped by evolution in a way that highly performing phenotypes have been selected [21]. Both FBA and cybernetic modelling have proved that this perspective can be used in different ways to simulate real biological behaviours. However, referring to the modelling categories defined in the introduction, FBA pertains to the group of *constraint-based* models, which are able to offer only a representation of the system at steady state. In FBA, the assumption is that the period in which the concentrations of biochemical species fluctuate can be neglected, while primary relevance is attributed to the condition in which the system is at equilibrium, i.e. when concentrations do not change in time any more. A further improvement of FBA is represented by **dynamic Flux Balance Analysis** (dFBA), proposed by Mahadevan and coworkers [19]. Here the extracellular space is modelled dynamically while inside the cell the steady-state assumption holds. In dFBA an ODE system is built to describe all extracellular events while the distribution of intracellular fluxes is calculated via a static optimization algorithm (SOA) or a dynamic optimization algorithm (DOA). For reasons of similarity with our approach we will focus specifically on the SOA algorithm, in which a biological objective function, like cellular growth rate or biomass, is maximized at each integration step of the ODEs. It is worth highlighting that dFBA always considers the intracellular system at steady state, so it neglects all the dynamic fluctuations of internal metabolites. In the field of mechanism based models, the CM [21] closely follows the teleonomic perspective, producing a long history of progressive refinements of their technique. Besides teleonomy, the CM approach also assumes that (i) the topology of the network is completely known, that (ii) metabolic enzymes are regulated at the gene expression level or via allosteric mech-

anisms, that (iii) some economic principles govern the way the cell implements these regulations, meaning that the cell, as if it were a rational investor, enhances those reactions that assure the highest yield in terms of the objective function. Finally CM assumes that (iv) accurate mathematical expressions with a complete list of kinetic parameters are available. These assumptions limit the good predictions of cybernetic models only to situations in which all parameters are available or where sufficient experimental data allow PE methods to be employed.

### 3. BACKGROUND

In this section we first introduce a brief overview of the (Stochastic) PN (SPN) formalism highlighting how it is possible to derive from an SPN model an ODE system that can be used to study the system’s behaviour. Subsequently, we will re-call the basis of the OPs and their solution techniques which will be used in the sequel of this paper.

**Stochastic Petri Nets.** PN and their extensions are graphical modelling formalisms which are becoming quite popular to build models of biological systems. The reason of this appeal is their capabilities of representing in a simple and intuitive manner many important features of these systems and thus of constructing models that are easy to understand also by non-mathematicians and non-computer scientists. PNs are bipartite directed graphs with two types of nodes called *places* and *transitions*. The *places*, graphically represented as circles, correspond to the system variables (e.g. enzymes and compounds), while the *transitions*, graphically represented as rectangles, encode the events (e.g. interactions among biochemical entities) which cause the system evolution. Arcs connecting places to transitions (and vice versa) express the relations between states and event occurrences. Places can contain tokens, graphically represented as black dots, that in the context of systems biology, often describe the number of molecules of the corresponding entities. An example of a PN model is shown in Fig. 1 which describes glycolysis in human red blood cells.

The state of a PN, called *marking*, is defined as the number of tokens in each place. An example of marking for the PN in Fig. 1 is showed in the third column of the Table in Fig. 2 . The system evolution is given by the occurrence of enabled transitions, where a transition is enabled if each input place contains a number of tokens greater or equal than a given threshold defined by the multiplicity of the corresponding input arc. A transition occurrence, called *firing*, removes a fixed number of tokens from its input places and adds a fixed number of tokens to its output places. The multiplicities of the input/output arcs determine the number of tokens involved by transition firings. The set of all the markings, that a net can reach through transition firings from an initial marking, is called the *Reachability Set* (RS). Instead, the behaviour of the net is encoded by means of the *Reachability Graph* (RG), a directed graph whose nodes are the markings of the RS and whose arcs are tagged with the labels of the transitions that cause the corresponding marking changes.

Temporal specifications must be introduced to model and study the temporal dynamics of a PN. Several timed extensions have been proposed in the literature [18]. In this paper we focus on Stochastic PN (SPN) [15], in which exponen-

tially distributed random delays (interpreted as durations of certain activities) are associated with transition firings. Thanks to this assumption the temporal behaviour of the system can be modelled with a random process governed by the so-called Chapman-Kolmogorov differential equations [5]. These equations correspond to the Master Chemical Equations [6] that are used to describe the behaviour of biological systems, thus making this formalism quite attractive for these types of applications. Specifically, the underlying stochastic process corresponds to a Continuous Time Markov Chain (CTMC) that can be represented as a graph isomorphic to the RG of the net. Formally an SPN can be defined as follows:

DEFINITION 1. *A stochastic Petri net system is a tuple*

$$\mathcal{N} = (P, T, I, O, \mathbf{m}_0, \lambda)$$

where:

- $P = \{p_i\}_{1 \leq i \leq n_p}$  is a finite and non empty set of places of cardinality  $n_p$ ;
- $T = \{t_i\}_{1 \leq i \leq n_t}$  is a finite, non empty set of transitions with cardinality  $n_t$  and such that  $P \cap T = \emptyset$ . All these transitions are *Timed transitions* which fire with a random delay characterized by a negative exponential probability distribution;
- $I, O : P \times T \rightarrow \mathbb{N}$  are the input, output functions that define the arcs of the net and that specify their multiplicities;
- $\mathbf{m}_0 : P \rightarrow \mathbb{N}$  is a multiset on P representing the initial marking; the notation  $m_0(p_i)$  specifies the initial marking of the place  $p_i$ ;
- $\lambda : T \rightarrow \mathbb{R}$  gives the firing intensities of the transitions.

The values assumed by the functions  $I$  and  $O$  can be collected in  $n_p \times n_t$  matrices (which we still call  $I$  and  $O$ ) and whose entries are  $I(p_i, t_j)$  and  $O(p_i, t_j)$ , respectively. By  $I(t)$  we denote the column of  $I$  corresponding to transition  $t$  (the same holds for  $O$ ). The set of input places of transition  $t$  (i.e. the preset of  $t$ ), denoted  $\bullet t$ , and the set of output places of  $t$  (i.e. postset of  $t$ ), denoted  $t^\bullet$ , are defined as follows:  $\bullet t = \{p \in P \mid I(p, t) \neq 0\}$ , and  $t^\bullet = \{p \in P \mid O(p, t) \neq 0\}$ .

**From SPN to ODE.** It often happens that, in case of very complex models, the underlying CTMC can not be derived or/and solved due to the well-known state space explosion problem. To cope with this difficulty, whenever the stochasticity of the modelled system can be neglected (e.g. due to huge number of molecules), the so-called deterministic approach can be exploited, assuming that the behavior of entities contained in a place of the net is described with an Ordinary Differential Equation (ODE) and that the whole model is specified with a system of ODEs, one for each place of the net. In the literature, different laws (e.g. Michaelis-Menten, Hill-equation, etc.) have been proposed to encode each reaction of the biological system into an ODE. Here we focus on the Mass Action (MA) law [20]<sup>1</sup> in which the ODEs describing the model have the following form:

$$\frac{dx_{p_i}(\nu)}{d\nu} = \sum_{j:O(p_i,t_j) \neq 0} O(p_i,t_j)\lambda(t_j) \prod_{h:I(p_h,t_j) \neq 0} x_{p_h}(\nu)^{I(p_h,t_j)} - \sum_{j:I(p_i,t_j) \neq 0} I(p_i,t_j)\lambda(t_j) \prod_{h:I(p_h,t_j) \neq 0} x_{p_h}(\nu)^{I(p_h,t_j)} \quad (1)$$

<sup>1</sup>Observe that this choice does not affect the generality of our approach that can be applied independently of the assumed law.

where  $x_{p_i}(\nu)$  represents the amount of the entity in place  $p_i$  at time  $\nu$  assuming that  $x_{p_i}(0)$  is defined through the initial marking of the net so that  $x_{p_i}(0) = m_0(p_i)$ .

For instance, considering the PN model in Fig.1 the behavior of place *GLC* is described by the following ODE equation assuming the MA law:

$$\begin{aligned} \frac{dx_{GLC}(\nu)}{d\nu} = & +\lambda(K_{F_1}) \cdot x_{HK} \cdot x_{GLC} \cdot x_{ATP} \quad (2) \\ & -\lambda(K_{R_1}) \cdot x_{HK} \cdot x_{G6P} \cdot x_{ADP} \end{aligned}$$

**Optimization problem.** In Mathematics, Computer Science, and Operations Research, optimization or mathematical programming consists of minimizing (or maximizing) a function by systematically choosing the values of its variables from a set of feasible possibilities properly exploiting analytical or numerical methods. In Systems Biology optimization is not a new concept since it has been already proposed to reconstruct gene regulatory networks, transcriptional regulatory networks, protein interaction networks, conditional specific sub-networks, and active pathways [10], and to perform FBA. Formally an optimization problem with inequality constraints can be defined as follows:

$$\begin{aligned} \underset{\mathbf{y}}{\text{minimize}} \quad & \mathcal{F}_{opt}(\mathbf{y}) \\ \text{subject to} \quad & \mathcal{G}_i(\mathbf{y}) \geq b_i, \quad 1 \leq i \leq l \\ & \mathcal{L}_j(\mathbf{y}) \leq c_j, \quad 1 \leq j \leq m \end{aligned}$$

where the vector  $\mathbf{y} = (y_1, \dots, y_n)$  is the *variable vector*, the function  $\mathcal{F}_{opt} : \mathbb{R}^n \rightarrow \mathbb{R}$  is the *objective function*, the functions  $\mathcal{G}_i(\mathbf{y}) : \mathbb{R}^n \rightarrow \mathbb{R}$  and  $\mathcal{L}_j(\mathbf{y}) : \mathbb{R}^n \rightarrow \mathbb{R}$  are *inequality constraint functions*, and the constants  $b_1, \dots, b_l, c_1, \dots, c_m$  are the *bounds* for the constraints. A vector  $\mathbf{y}^*$ , called *optimal*, is the solution of the OP if, among all vectors that satisfy the constraints, it is that which yields the smallest (largest) value of the optimization function:  $\forall \mathbf{z}$  s.t.  $\mathcal{G}_i(\mathbf{z}) \geq b_i, \dots, \mathcal{L}_j(\mathbf{z}) \leq c_j$  we have that  $\mathcal{F}_{opt}(\mathbf{z}) \geq \mathcal{F}_{opt}(\mathbf{y}^*)$ .

We recall that an OP is called a *linear program* if the objective and constraint functions are linear and *non-linear* otherwise. As shown in the next sections of this paper, we will focus on non-linear programs in which constraints can be non-linear as well. To solve this type of OPs, several algorithms have been proposed in the literature, and the reader can find a complete survey of these methods in [11].

## 4. OUR APPROACH

Before describing our approach in details, we introduce the biological considerations which motivated our proposal.

First, in our method we assume that both the topology and the kinetic parameters of the network are specified with different level of detail, depending on the biochemical event we are observing. Since in biochemical systems events are generally represented as biochemical reactions, we decide to divide all the reactions into two classes: *determined* reactions and *undetermined* reactions.

We assume that for determined reactions the mechanism of their biochemical interaction has been properly analysed, so that they can be clearly characterized and modelled.

Instead, regarding undetermined reactions, we assume that in this part of the network the topology is only partially known, so that while all the reagents, all the products and some modifiers are specified, we do not exclude that other modifiers could be present.

For instance, this assumption can be motivated by many studies that showed how many cells of different organisms own different isoforms of the same enzyme and that every isoform has a specific affinity for a group of molecular regulators [16]. Thus, in a specific cell, the resulting kinetic behaviour of a reaction is a direct consequence of the proportion at which these isoforms are present [16] [2]. In some situations this proportion may hardly be defined, and, even if it is known, the list of modifiers for secondary isoforms is not always clear.

Moreover, for specific conditions, like cancer, cells may present genetic mutations and quantitative alterations or isoform switches that change the affinity between enzymes and their regulators, thus affecting the overall enzymatic dynamics. Assuming the molecular interactions of an enzyme are unclear necessarily implies that its activity is not expressible with a precise mathematical formulation.

Finally, as anticipated in Section 2, we assume that an objective function with a biological meaning can be profitably used to formulate an OP and to reproduce an actual biological behaviour. From our point of view, this biological objective can be interpreted in different ways. For instance, it can be related to its definition in FBA or CM and formulated similarly. In FBA, according to the teleonomic perspective, the objective function describes an objective that we believe is physiologically relevant for the cell, like the maximisation of biomass. This is used to predict a physiological behaviour of the cell. Inspired by other successful applications of FBA, also in our approach the objective function can express a non-physiological goal of the cell, like the maximisation of ATP production, and used to identify some property of the network, or, it may be used to impose an engineering (e.g. maximisation of the production of a particular aminoacid) or therapeutic (e.g. minimisation of some reaction fluxes vital for cancer cells) objective to a metabolic system. More in general, the objective function is intended to represent any relevant biological behaviour that has been experimentally measured or that has to be achieved. Our approach thus tries to investigate if and how this specific biological phenotype can be reproduced in a biochemical system where the topology and the parametrization are just partially known.

In details we propose a method that exploits iterative optimizations: at each simulation step, an objective function with the aforementioned biological meaning allows us to estimate the activities of undetermined reactions, and thus to obtain a complete description of system behaviour.

To facilitate the construction of the model we propose a new graphical formalism based on PN, which allows to automatically translate the model into its mathematical representation, consisting of the ODEs system and the Optimization Problem (OP).

According to this we decide to present our approach firstly introducing this new graphical formalism, and then providing its automatic translation into a ODE system in which indeterminate transitions are tackled through an OP. For this purpose we use the model of Fig.1 as a “running example” that we comment in the rest of the paper to discuss the features of this new modelling formalism.

**SPN with Indetermination.** The formal definition of a new PN extension called Stochastic Petri Net with Indetermination (SPNI) is the following:

DEFINITION 2. A stochastic Petri net with indetermina-  
tion is a tuple

$$\mathcal{N} = (P, T, I, O, \mathbf{m}_0, \lambda_n, \Lambda_u, \mathcal{F}_{opt}^N)$$

where:

- $P, I, O, \mathbf{m}_0$  are defined as in the SPN formalism;
- $T = T_n \cup T_u$  is a finite, non empty set of timed transitions with  $T_n \cap T_u = \emptyset$ .  $T_n$  is the set of the determined transitions, while  $T_u$  is the set of undetermined transitions.
- $\lambda_n : T_n \rightarrow \mathbb{R}$  gives the firing intensity of  $T_n$  transitions.
- $\Lambda_u : T_u \rightarrow \mathbb{R}^2$  gives the range of variation of the flux of  $T_u$  transitions.
- $\mathcal{F}_{opt}^N : T \times P \rightarrow \mathbb{R}$  is an objective function whose terms are represented by place markings and transition firing intensities.

We use the notation  $\Lambda_u^L(t)$  (resp.  $\Lambda_u^H(t)$ ) to denote the lower (resp. higher) bound values of the firing intensity interval associated with a  $t \in T_u$ ;  $\Lambda_u(t)$  then represents a possible firing intensity of the (undetermined) transition  $t$  compatible with its specified lower and higher bounds. For instance in the gray boxes of Fig.1 are represented the transitions affected by a level of indetermina-  
(i.e.  $K_{f1}, K_{r1}, K_{f3}, K_{r3}, K_{f12}, k_{r12}$ ).

#### From SPNI to ODE and OP.

Due to the indetermina-  
tion associated with the  $T_u$  transi-  
tions, it is not possible to directly use Eq. 1 to represent the deterministic behavior of an SPNI model. We can however re-write Eq. 1 as follows:

$$\begin{aligned} \frac{dx_{p_i}(\nu)}{d\nu} = & \sum_{j:O(p_i,t_j) \neq 0} O(p_i,t_j) \mathcal{M}_{t_j}(\nu) \prod_{h:I(p_h,t_j) \neq 0} x_{p_h}(\nu)^{I(p_h,t_j)} \\ & - \sum_{j:I(p_i,t_j) \neq 0} I(p_i,t_j) \mathcal{M}_{t_j}(\nu) \prod_{h:I(p_h,t_j) \neq 0} x_{p_h}(\nu)^{I(p_h,t_j)} \end{aligned} \quad (3)$$

where  $\mathcal{M}$  is a function defined in the following way:

$$\mathcal{M}_t(\nu) = \begin{cases} \lambda_n(t) & \text{if } t \in T_n \\ y_t(\nu) & \text{otherwise} \end{cases} \quad (4)$$

The parameter  $y_t(\nu)$  encodes the indetermina-  
tion associated with the undetermined transition  $t$  at time  $\nu$  and must be properly estimated to solve the ODE system.

As we already pointed out, the undetermined transitions are part of the model either because their exact specification is not relevant with respect to the goals of the analysis carried on with the SPNI or because they are too difficult to identify in a precise manner. Independently of the context of the modelling experiment, it is usually the case that we want to minimize (or maximize) certain measures defined on the portion of the state of the system that is not directly affected by undetermined transitions. These measures, that may assume complex definitions, become the optimization functions that we use to study these models. To cope with this problem we thus propose to proceed in two steps where we first reduce the indetermina-  
tion of the model solving an optimisation problem in which the firing intensity of  $T_u$  transitions are estimated according to a user-defined objective function. Subsequently, we use these estimated values in the ODE system which thus becomes properly defined and ready to be solved.

Given an SPNI model, the corresponding OP, whose solution will be used to estimate the firing intensity values of the  $T_u$  transitions, is derived using the following definition.

DEFINITION 3. The OP derived by the SPNI is a tuple

$$\mathcal{O} = (\mathbf{y}_\nu, \mathcal{F}_{opt}, \mathcal{G}, \mathcal{L})$$

where:

- $\mathbf{y}_\nu$  represents the optimizing values of undetermined transitions at time  $\nu$ , i.e.  $\forall t \in T_u \Rightarrow y_t(\nu) \in \mathbf{y}_\nu$ ;
- $\mathcal{F}_{opt} = \mathcal{F}_{opt}^N$ ;
- $\mathcal{G}$  is defined by

$$\forall t \in T_u \Rightarrow y_t(\nu) \prod_{h:I(p_h,t) \neq 0} x_{p_h}(\nu)^{I(p_h,t)} \geq \Lambda_u^L(t)$$

- $\mathcal{L}$  is defined by

$$\forall t \in T_u \Rightarrow y_t(\nu) \prod_{h:I(p_h,t_j) \neq 0} x_{p_h}(\nu)^{I(p_h,t_j)} \leq \Lambda_u^U(t)$$

For instance, considering the SPNI in Fig. 1 the vector  $\mathbf{y}(\nu)$  has size six and represents the optimal values of the firing rates of transitions  $K_{f1}, K_{r1}, K_{f3}, K_{r3}, K_{f12}$ , and  $K_{r12}$ . An example of objective function could be the maximization of the Lactate (LAC) as described in our case study in Section 5. Moreover in our example  $\mathcal{G}(K_{f1})$  is  $y_{K_{f1}} \cdot y_{HK} \cdot x_{G6P} \cdot x_{ADP} \geq 8.0E^5$  and  $\mathcal{L}(K_{f1})$  is  $y_{K_{f1}} \cdot x_{HK} \cdot x_{G6P} \cdot x_{ADP} \leq 9.0E^4$  with limit values chosen as explained in Section 5

**How to compute the model behavior.** The numerical integration of Eq. 3 provides the behavior of the model  $\mathbf{x}(\nu + \tau)$  at time  $\nu + \tau$ , in terms of the behavior  $\mathbf{x}(\nu)$  computed at time  $\nu$  and of a set of parameters deriving from the structure of the SPNI ( $I$ , and  $O$ ), the firing intensities of the definite transitions of the net ( $\lambda_n$ ) and of the firing intensities of the undetermined transitions estimated at time  $\nu$  and collectively represented as  $\mathbf{y}(\nu)$ . The values of  $\mathbf{x}(\nu + \tau)$  are thus the results of the evaluation of a function whose input parameters are represented by a tuple  $B(\nu) = (B, B_u(\nu))$  where  $B = (I, O, \lambda_n)$  and  $B_u(\nu) = (\mathbf{x}(\nu), \mathbf{y}(\nu))$ <sup>2</sup>. The integration step  $s$  identifies the time points  $\nu_i = i * \tau$  where the evaluation of the model behavior is of interest.

Role of the estimation phase of our method is that of finding a set  $\mathbf{y}(\nu)$  that, being compatible with the constraints of the SPNI model ( $\Lambda_u$ ), minimizes the objective function at time  $\nu + s$ . The optimization phase identifies a number  $K$  of initial conditions, that we denote with  $B_u^{[k]}(\nu), k = 1, \dots, K$ , consisting of the behavior of the model computed at time  $\nu$  and of  $K$  random points within the space of firing intensities of the undetermined transitions identified by the constraints  $\Lambda$ . From each of these configurations the method numerically integrates the system of ODEs up to time  $\nu + s$  to derive  $\mathbf{y}(\nu)$ . Letting  $B^{[k]}(\nu) = (B, B_u^{[k]}(\nu))$ , with  $B_u^{[k]}(\nu) = (\mathbf{x}(\nu), \mathbf{y}^{[k]}(\nu))$ , the solutions obtained from the integration of the ODEs with parameters  $B^{[k]}(\nu), k = 0, 1, \dots, K$  and up to time  $\nu + s$  are compared to identify the choice of  $B^{[k]}(\nu)$  which provides the best evaluation of the objective function, thus identifying  $\mathbf{y}^{[k]}(\nu + s) = \mathbf{y}(\nu)$ . Crucial in this optimization step is that the numerical integration of the ODEs is performed with a method capable of identifying an integration step  $h$  small enough to allow a precise solution of the

<sup>2</sup>In the sequel of the paper we will indifferently use  $\mathbf{y}_t(\nu)$  or  $\Lambda_u(t, \nu)$  to represent the undetermined parameters of our models as provided by the optimization problem at time  $\nu$ .

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**Algorithm 1** Algorithm to solve ODE system with Indetermination

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1: function SOLVEODEI(ODEI,  $\mathcal{G}$ ,  $\mathcal{L}$ ,  $\mathcal{F}_{opt}$ ,  $\tau$ , FinalTime)
2:    $\nu = 0.0$ ;
3:   ODEI.Init(Value);
4:   while ( $\nu \leq$  FinalTime) do
5:     print(time, Value);
6:      $Rate_{T_u} = SolveOpt(Value, ODEI, \nu + \tau, \mathcal{G}, \mathcal{L}, \mathcal{F}_{opt})$ ;
7:      $Value = ODEI.SolverODE(Value, \nu + \tau, Rate_{T_u})$ ;
8:      $\nu += \tau$ ;
9:   end while
10: end function

```

---

ODEs during these “tentative” evaluations that are used to select the firing intensities of the undetermined transitions.

In general, this whole method is repeated for each time point  $\nu_i$  starting from  $\nu_0 = 0$ . However, solving the OP for each value of  $\nu_i$  can be excessively costly and we can thus reduce this computational effort by identifying a time interval  $\rho$  that is a multiple of  $\tau$  and that determines the time points where the optimization is requested. By doing so, if we set  $\rho = m \cdot \tau$ , we assume that for  $m - 1$  intermediate evaluation steps the values of  $\Lambda_u(\nu)$  (i.e.  $\mathbf{y}(\nu)$ ) remain constant and an approximation is introduced.

Having discussed how to derive from an SPNI model (i) an ODE system with indetermination (see Eq.3), and (ii) an OP (see Def.3), we can devise an algorithm which combines them to derive the model behavior.

The pseudo-code of this algorithm is shown in Alg. 1. It takes as input the ODE system with indetermination (i.e. *ODEI*), the OP (i.e. described by functions  $\mathcal{G}$ ,  $\mathcal{L}$  and  $\mathcal{F}_{opt}$ ), the step size used for the optimization schema (i.e.  $\tau$ ), and the final time (i.e. *FinalTime*). The output of the algorithm is represented by the values generated for each system entity at different time points (i.e.  $\nu_i$ ). In details, the method *Init()* at line 3 initializes the vector *Value* encoding the initial values assumed for all the entities of the model. Then, the code from line 6 to line 8 is repeated until the time horizon is not reached. In each iteration the function *print()* is called to print the current values of the system entities. Subsequently, the function *SolveOpt()* solves the optimization and returns an estimated value for each  $T_u$  transition. It takes in input the current values of the entities (i.e. *Value*), the final time in which the objective function will be evaluated (i.e.  $\nu + \tau$ ), the ODE system (i.e. *ODE*), and a set of functions encoding the OP (i.e.  $\mathcal{G}$ ,  $\mathcal{L}$  and  $\mathcal{F}_{opt}$ ). The functions  $\mathcal{G}$  and  $\mathcal{L}$  are used by the optimization solver to test if a new vector  $\mathbf{y}$ , randomly generated according to the parameter constrains, is a feasible solution. Indeed the functions  $\mathcal{G}$  and  $\mathcal{L}$  verify if  $\mathbf{y}$  satisfies the inequality constraints. The function  $\mathcal{F}_{opt}$  is instead called by the optimization solver to compute the value of the objective function associated with a feasible vector  $\mathbf{y}$ . This function, takes as input the vector  $\mathbf{y}$ , the current values of the entities (i.e. *Value*), the ODE system (i.e. *ODE*), and the final time in which the objective function must be evaluated (i.e.  $\nu + \tau$ ). First it computes the quantities values at  $\nu + \tau$  assuming the missing rates to be equal to  $\mathbf{y}$ . Then, the computed values are used to evaluate the objective function, whose derived value is returned. When the optimization step is terminated returning the rate estimations, the ODE system with indetermination (i.e. *ODEI*) can be solved (i.e. method *SolveODE()*), and the vector *Value* is updated with the new computed values.

## 5. EXPERIMENTAL RESULTS AND DISCUSSION

In this section we report a case study to illustrate how our approach can be used to investigate the metabolic behaviour of cancer cells. The model represents the glycolytic pathway in a generic human cell. It is inspired by the model presented in [9], which describes glycolysis in human red blood cells. In physiological conditions glycolysis leads to a consumption of Glucose (GLC) and a progressive production of Pyruvate (PYR) and energy, in the form of Adenosine Triphosphate (ATP). Then, in the presence of oxygen, PYR is metabolised by other pathways to generate the majority of the energy consumed by the cell. In absence of oxygen, PYR is converted to LAC without further energetic yields. Differently from normal cells, cancer cells exhibit an enhancement of glycolysis and production of LAC even in the presence of oxygen, Warburg Effect [7]. This phenomenon inspired the experiment discussed below. The model is characterized by fourteen metabolic reactions, the related equations are reported in the first column of the Table in Fig. 2, and it can be graphically described by the SPN model in Fig. 1 where place names are chosen to recall the corresponding biological compounds.

**Our implementation.** The experiments have been performed using a prototype implementation of the proposed method integrated in the *GreatSPN* framework [1]. With this extension, *GreatSPN* allows the generation of an ODE system with/without indetermination from an SPNI model, and then the *R environment* is used to compute its solutions. In particular the *deSolve* library is used to solve the ODE system, while *GenSA* library to solve the OP.

As already pointed out, in cancer cells the metabolism is characterised by the Warburg Effect [7]. Moreover, it has been recently shown that cancer cells preferentially rely on specific isoforms of their metabolic enzymes. In particular, it seems that isoforms of **Hexokinase** (HK), **Phosphofruktokinase** (PFK) and **Pyruvate Kinase** (PK) may have an eminent role in the deregulation of cancer metabolism [13]. Despite these discoveries, it is still complicated to characterize the *in vivo* kinetics of these isoforms. Conditioned by these constraints, we chose to set the reactions involving HK, PFK and PK as undetermined transitions, i.e deficient of a complete list of regulators, of its mathematical expression and its parametrization. Our approach is here used as an attempt to acquire a deeper understanding of these metabolic dynamics. The idea is to use an objective function that encodes the Warburg Effect in every type of cancer at every possible tumour stage. For this reason the firing intensities  $K_{f_1}$ ,  $K_{r_1}$ ,  $K_{f_3}$ ,  $K_{r_3}$ ,  $K_{f_{12}}$  and  $K_{r_{12}}$  were allowed to vary in a wide range that agrees with biological knowledge. This was formalized as the maximization of LAC production at every integration step.

Fig. 3 shows the evolution of LAC over time. The black line represents the results of the model of a normal cell, where all parameters are well characterized. Blue triangles show the time evolution of LAC when uncertainty is applied. It can be noticed that this objective function is able to drive the system to accelerate LAC production. Even if the difference might not seem large enough to represent the Warburg Effect, we point out that these diagrams show the behaviour of our model for a very short time interval. Fig. 4 shows the

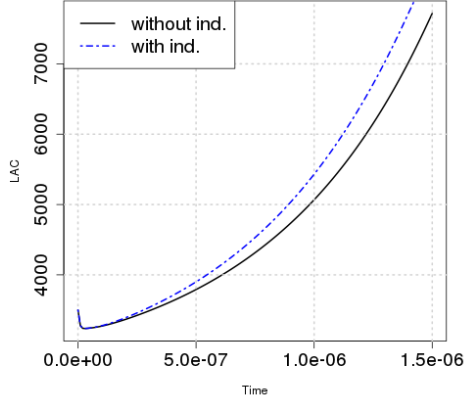


Figure 3: Behaviours of place *LAC*

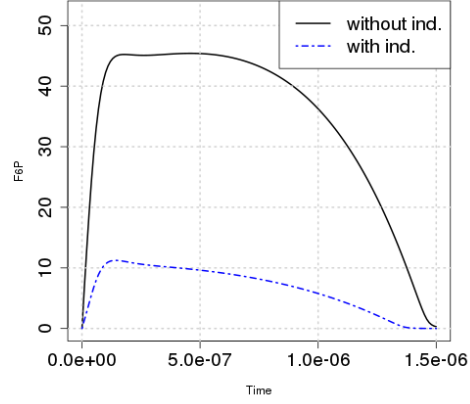


Figure 4: Behaviours of place *F6P*

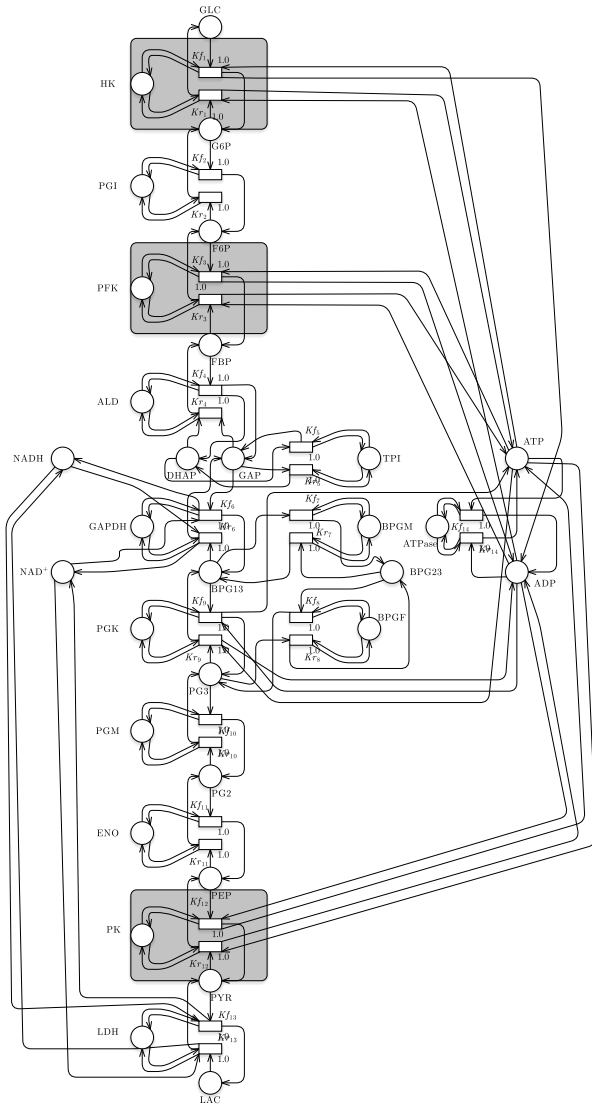


Figure 1: Case study: Glycolysis in *Homo Sapiens*.

Reactions	Rate Equations	Initial Marking
$HK + GLC + ATP \xrightarrow{K_{f1}} HK + G6P + ADP$	$K_{f1} = 6.48E+05, K_{r1} = 4.9E+03$	$GLC_0 = 5E+03, HK_0 = 4$
$PGI + G6P + ATP \xrightarrow{K_{f2}} PGI + F6P$	$K_{f2} = 1.15E+03, K_{r2} = 2.68E+03$	$G6P_0 = 0, PGI_0 = 8E+03$
$PFK + F6P + ATP \xrightarrow{K_{f3}} PFK + FBP + ADP$	$K_{f3} = 1E+09, K_{r3} = 8.47E+04$	$F6P_0 = 0, PFK_0 = 10$
$ALD + FBP \xrightarrow{K_{f4}} ALD + DHAP + GAP$	$K_{f4} = 1.46E+02, K_{r4} = 1.18E+00$	$FBP_0 = 0, ALD_0 = 2E+04, DHAP_0 = 0$
$TPI + GAP \xrightarrow{K_{f5}} TPI + DHAP$	$K_{f5} = 7.93E+00, K_{r5} = 4.53E-01$	$TPI_0 = 2E+06, GAP_0 = 0$
$GAPDH + GAP + NAD^+ \xrightarrow{K_{f6}} GAPDH + BPG13 + NADH$	$K_{f6} = 1.42E+05, K_{r6} = 5.28E+06$	$GAPDH_0 = 4E+02, NAD_0^+ = 1E+04, NADH_0 = 0$
$BPGM + BPG13 \xrightarrow{K_{f7}} BPGM + BPG23$	$K_{f7} = 1E+08, K_{r7} = 1E+05$	$BPGM_0 = 5, BPG13_0 = 0$
$BPGF + BPG23 \xrightarrow{K_{f8}} BPGF + PG3$	$K_{f8} = 6.84E+02, K_{r8} = 1E-09$	$BPGF_0 = 1E+02, BPG23_0 = 0$
$PGK + BPG13 + ADP \xrightarrow{K_{f9}} PGK + PG3 + ATP$	$K_{f9} = 2.61E+04, K_{r9} = 1.45E+01$	$PGK_0 = 1E+02$
$PGM + PG3 \xrightarrow{K_{f10}} PGM + PG2$	$K_{f10} = 5.38E+01, K_{r10} = 7.92E+00$	$PGM_0 = 9E+05, PG3_0 = 0$
$ENO + PG2 \xrightarrow{K_{f11}} ENO + PEP$	$K_{f11} = 5.82E+02, K_{r11} = 3.44E+02$	$ENO_0 = 1E+05, PG2_0 = 0$
$PK + PEP + ADP \xrightarrow{K_{f12}} PK + PYR + ATP$	$K_{f12} = 5.17E+02, K_{r12} = 5.17E-01$	$PEP_0 = 0, PK_0 = 9E+03$
$LDH + PYR + NADH \xrightarrow{K_{f13}} LDH + LAC + NAD^+$	$K_{f13} = 1.04E+03, K_{r13} = 2.34E+00$	$PYR_0 = 1E+02, LDH_0 = 4E+02, LAC_0 = 3.5E+03$
$ATPase + ATP \xrightarrow{K_{f14}} ATPase + ADP$	$K_{f14} = 9.74E-01, K_{r14} = 9.74E-04$	$ATPase_0 = 1E+02, ATP_0 = 7E+02, ADP_0 = 5E+02$

Figure 2: Table: Reactions, Equations and Initial marking of glycolysis in *Homo Sapiens*.

rapid accumulation of Fructose 6-Phosphate (F6P) in the normal cell model compared to a more balanced production-consumption dynamics in the cancer model. F6P, a high glycolytic intermediate, increases as a direct consequence of GLC degradation and is later processed by PFK. It is then significant to see which parameters the optimizer independently chooses to tune in order to maximize its objective function. While parameter values of HK and PK did not vary markedly if compared to the normal cell model (data not shown), the highest difference regarded PFK. Many articles as [16] have demonstrated that PFK kinetics is highly non-linear and depends on many allosteric interactions. Our results seem to reinforce the Mulukutla's thesis [16] that the regulation of PFK activity has a crucial impact on the glycolytic flux and may be relevant to explain metabolic alterations in cancer.

**Discussion.** It is important to observe that our approach differently by dFBA and CM allows us to define objective functions that may mimic the behaviours experimentally observed, as the LAC maximisation. Moreover, our proposed approach does not assume the steady state of the intracellular metabolites as in dFBA, and it does not require a full understanding of model kinetics as in CM. Finally, unlike PE it is not limited by the lack of experimental data, as

normally happens in cancer cell scenarios.

## 6. CONCLUSIONS

In this work we proposed a new approach to deal with the biochemical models with lacking kinetic information. As discussed in Section 5 it may be useful for different applications, although additional experiments will provide a further validation and will show its actual relevance. For the future we intend to apply our approach to study different networks, either in microorganisms, in single human cells or in population of cells. Because the performances of our approach are highly dependent on the optimization algorithms we plan to test different algorithms comparing their performance in terms of memory saving and execution time. Great efforts will be also spent to investigate which available biological data can be effectively used to adapt the model to the specific situation under study. Transcriptomics, proteomics and metabolomics profiles, as well as reaction fluxes and thermodynamic information are all possible candidates. These data will provide a fundamental support to define the initial conditions of the model and to set the constraints for the optimization. Finally, we will investigate the possibility of extending our approach to grab stochastic behaviours, so that the model will be approximated by a system of stochastic differential equations.

## 7. ADDITIONAL AUTHORS

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